NRC FORM 591M PART (08-2010) 10 CFR 2.201		DECTION DED	U.S NUCLEAR REGULATORY COMMISSION ORT AND COMPLIANCE INSPECTION			
	SAFETT INS	FEO HON REF	OK! AND COMP	LIANCE INSPEC	HON	
1. LICENSEE/LOCATION INSPECTED:			2, NRC/REGIONAL OFFICE			
West Branch Regional Medical Center 2463 South M-30 West Branch, MI 48661			U.S. Nuclear Regulatory Commission Region III 2443 Warrenville Road, Suite 210 Lisle, Illinois 60532-4351			
REPORT NUMBER(S) 2011-01						
3. DOCKET NUMBER 030-173		4 LICENSEE NUMB 21-18	ER(S) 3892-01	5 DATE(S) OF INSI June 22 and J		
LICENSEE:						
compliance with the	Nuclear Regulator of selective exami	y Commission (NRC inations of procedure	ted under your license a i) rules and regulations a as and representative re as follows:	and the conditions of vi	our license. The	
1. Based on the inspection findings, no violations were identified.						
2. Previous	violation(s) closed.					
3. The violation(s), specifically described to you by the inspector as non-cited violations, are not being cited because they were self-identified non-repetitive, and corrective action was or is being taken, and the remaining criteria in the NRC Enforcement Policy, NUREG-1600, to exercise discretion, were satisfied						
Non-cited violation(s) were discussed involving the following requirement(s):						
4. During this inspection certain of your activities, as described below and/or attached, were in violation of NRC requirements and are being cited. This form is a NOTICE OF VIOLATION, which may be subject to posting in accordance with 10 CFR 19.11 Title 10 Code of Federal Regulations (CFR) 35.67(g) requires that a licensee in possession of sealed sources or brachytherapy sources, except for gamma stereotactic radiosurgery sources, shall conduct a semi-annual physical inventory of all such sources in its possession. The licensee shall retain each inventory record in accordance with § 35.2067(b). Continued on Part 2						
		Statement o	f Corrective Actions	**************************************		
I hereby state that, violations identific 2.201 (corrective st achieved). I unders	within 30 days, th d. This statement teps alre⊔dy taker stand that no furth		ed by me to the inspect ns is made in accordar which will be taken, de se to NRC will be requir	tor will be taken to co nce with the requirem to when full complian red, unless specifical	rrect the lents of 10 CFR nce will be ly requested.	
Title	Printed	Name	Signat	uro /	Date	
LICENSEE'S REPRESENTATIVE				ombah	9-6-2011	
NRC INSPECTOR	Robert F	P. Hays	2600	a de la companya della companya della companya de la companya della companya dell	8/19/2011	
Branch Chief	Tamara E.		Taurara	Slovens	9/8/11	
NRC FORM 591M PAI	RT 1(06-2010)		Town do	/	11010	

NRC FORM 591M PART 2 (06-2010) 10 CFR 2:201 SAF	ETY INSPECTION RE	U.S NUCLEAR REGULATORY COMMISSION PORT AND COMPLIANCE INSPECTION		
1 LICENSEF/LOCATION INSPECTED: West Branch Regional Me 2463 South M-30 West Branch, MI 48661 REPORT NUMBER(S) 2011-01	edical Center	2. NRC/REGIONAL OFFICE U.S. Nuclear Regulatory Commission Region III 2443 Warrenville Road, Suite 210 Lisle, Illinois 60532-4351		
3. DOCKET NUMBER(S)	4. LICENSEE NUMBE	(R(G)	5. DATE(S) OF INSPECTION	
030-17321	21	-18892-01	June 22 and July 28, 2011	

(Continued)

Contrary to the above, the licensee has not conducted a semi-annual inventory of brachytherapy sources as required. Specifically, the licensee has not conducted a semi-annual inventory of the brachytherapy seeds used for prostate implants since January 20, 2008. The licensee has maintained a record of unused brachytherapy seeds placed in storage, but failed to conduct a semi-annual inventory of those seeds in storage to ensure that no seeds have become lost or unaccounted for.

This is a Severity Level IV violation (Section 6.3).

The licensee's medical physicist believed that an annual inventory was required rather than a semi-annual inventory. The licensee's corrective actions will be to conduct an inventory of the brachytherapy seeds with the next 30 days and will continue to conduct an inventory of the brachytherapy seeds semi-annually as required.

NRC FORM 591M PART 2(08-2010)

NRC FORM 591 M PART 3		U.S. NUCLEAR REGULATORY COMMISSION					
(06-2010) 10 CFR 2.201 SAFETY	NSPECTION R	Docket File Information EPORT AND COMPLIANCE IN	SPECTION				
1. LICENSEE West Branch Regional Med 2463 South M-30 West Branch, MI 48661 REPORT NUMBER(S) 2011-01	ical Center	2. NRC/REGIONAL OFFICE U.S. Nuclear Regulatory Commission Region III 2443 Warrenville Road, Suite 210 Lisle, Illinois 60532-4351					
3. DOCKET NUMBER(S) 03017321	4. LICENSE NUM	MBER(S) 21-18892-01	5. DATE(S) OF INSPECTION June 22, 2011				
6. INSPECTION PROCEDURES	7. INSPECTION		Cuito EE, EU11				
87131 (10/24/02)		03.01-03.07					
SUPPLEMENTAL INSPECTION INFORMATION							
1.PROGRAM 2. PRIORITY 3			4. TELEPHONE NUMBER 989-345-3660				
X Main Office Inspectio ☐ Field Office Inspectio ☐ Temporary Job Site In	on	Next Inspection	Date: <u>June 2014</u>				

PROGRAM SCOPE

The licensee was authorized for two locations in West Branch, Michigan, with authorization by the license at the main campus, 2463 S. M-30, to use any byproduct materials for diagnostic and therapeutic medical procedures under 10 CFR 35.100, 35.200, 35.300, and 35.400. The second location, West Branch Regional Cancer Center, 2431 S. M-30, is authorized to use any byproduct materials for diagnostic medical procedures under 10 CFR 35.100 and 35.200, excluding PET isotopes and Xe-133. Cardiac stress testing at the Cancer Center was inspected during the prior inspection and was not reviewed during the current inspection.

The licensee's main Nuclear Medicine Department routinely conducts a daily average of 4-6 patient studies with a staff of 2 nuclear medicine technologists. Iodine-131 procedures requiring a written directive average one or two procedures per year. One administration of Sr-89 has been performed since the previous inspection. The licensee receives licensed material as unit doses and bulk pertechnetate from an area local nuclear pharmacy as needed.

The licensee periodically performs iodine-125 seed implant procedures and averages 3 cases per year. Implant records are maintained in the dosimetry files in the Cancer Center for review. Unused seeds are stored for decay in the main campus hot lab.

Performance Observations

During the inspection, the licensee's NMT staff demonstrated/discussed: (1) survey instruments and required surveys; (2) package receipt and check-in procedures; (3) wipe test counting; (4) unit dose and safe handling procedures; (5) I-131 procedures and written directives; (6) waste handling; (7) sealed source inventories and leak tests; (8) security and storage of licensed material; (9) radiation safety program audit results; (10) dosimetry for CY 2009: 472mr-DDE; 1830mr-finger; and CY 2010: 425mr-DDE; 1380mr-finger. The inspector performed independent and confirmatory radiation measurements, which indicated results consistent with licensee survey records and postings. A SL IV violation of 10 CFR 35.76(g) was identified for a failure to perform semi-annual inventories of brachytherapy seeds.